- 1 -

## METHOD OF ASSAYING PYRROLE-CONTAINING BIOLOGICAL COMPOUNDS

1	Cross-Reference to Related Application
2	
3	This application is a divisional of co-pending
4	U.S. Application No. 09/970,328, filed October 2,
5	2001, which is a continuation-in-part of U.S.
6	Application No. 09/679,141, filed October 3, 2000
7	(now abandoned), the disclosures of which are
8	incorporated herein by reference.
9	
10	BACKGROUND OF THE INVENTION
11	Field of the Invention
12	This invention relates to methods of assaying
13	pyrrole-containing biological compounds and chemical
14	compositions that can be used in such methods. More
15	specifically, it relates to a method for detecting
16	pyrrole-containing molecules that are markets of
17	particular disease states.
18	
19	Description of Related Art
20	Erlich's reagent, or p-dimethylaminobenzaldehyde
21	(1), is a molecule that can react with pyrroles and
22	indoles to form a chromogenic compound.

2 See G. Lombard and V. Dowell, J. Clin. Microbiol.

3 (1983) 18:609-613. The mechanism of action is

4 typically described as an electrophilic attack on the

5  $\alpha$ -carbon atom of a pyrrole. This attack forms a

6 highly conjugated cation that absorbs light in the

7 visible spectrum. Such a mechanism is graphically

8 represented in Scheme A above.

9

10 The reaction of Ehrlich's reagent with certain

11 compounds has been discussed. For instance, Iyer

12 reported a pyrrole is formed when LGE2 is reacted

13 with proteins. See Iyer et al., J. Org. Chem. (1994)

14 59:6038-6043. When the pyrrole was contacted with

15 Ehrlich's reagent in the presence of BF3 OEt2, a blue-

16 green chromophore was produced. The chromophore was

17 identified as a pyrrolic electrophilic substitution

18 product.

1	Lombard reported the reaction between Ehrlich's
2	reagent and bacterially derived indoles. See G.
3	Lombard and V. Dowell, J. Clin. Microbiol. (1983)
4	18:609-613. The sensitivity of the reagent was
5 .	compared to two other indole detecting compounds:
6	Kovac's reagent and DMCA. Ehrlich's reagent was
7	reported to be 10 times less sensitive than DMCA and
8	10 times more sensitive than Kovac's reagent in
9	detecting indole.
10	
11	While Ehrlich's reagent has been used to roughly
12	detect the presence of pyrroles or indoles in a
13	targeted material, improved compositions and methods
14	for detecting such heterocycles are desirable,
15	especially methods that provide for detecting
16	pyrrole-containing molecules that are markers of
17	particular disease states.
18	
19	SUMMARY OF THE INVENTION
20	
21	The present invention provides methods of assaying
22	pyrrole-containing biological compounds.
23	In one case the method involves:
24	<ol> <li>contacting the biological compound with either:</li> </ol>
25	a) an optionally labelled derivatizing agent
26	(bound to or able to bind to a solid
27	support), wherein the derivatizing agent
28	forms a reaction product with the
29	biological compound (preferably via
30	covalent attachment thereto), followed by

	1		exposure to a detectable molecule which
	2		forms a complex with the reaction product;
	3		or
	4	b)	an optionally labelled derivatizing agent
	5		not bound to a solid support, wherein the
	6		derivatizing agent forms a reaction product
	7		with the biological compound (preferably
	8		via covalent attachment thereto), followed
	9		by exposure to a binding agent specific to
	10		the biological compound in the reaction
	11		product, said binding agent being bound to
	12		a solid support; or
	13	c)	a binding agent bound to a solid support,
	14		said binding agent being specific to the
	15		biological compound and forming a complex
	16		therewith, followed by exposure to an
•	17		optionally labelled, derivatizing agent
	18		which forms a reaction product with the
	19		biological compound moiety of said complex
	20		(preferably via covalent attachment
	21		thereto); and
	22 ·		·
	23 2)	dete	ermining the amount of bound biological
	24	comp	pound by detecting the detectable molecule,
	25	or l	by determining the amount of free or bond
	26	bine	ding agent or by measuring the amount of
	27	labe	el present.
	28		

1 Preferably, the method of assaying pyrrole-containing

- 2 biological compounds is Method 1, described in part
- 3 a) above. Method 1 involves the following steps:

5 1) contacting a biological compound with a
6 derivatizing agent of the following structure in
7 the bound form;

wherein R<sup>1</sup> is an alkyl group, R<sup>2</sup> is an alkyl group, A is a linking group and B is a solid support, and wherein the contact induces formation of a reaction product, and wherein the reaction product comprises the covalent attachment of the biological compound to the derivatizing agent; followed by contacting the reaction product with a detectable molecule, wherein the contact induces specific binding of the detectable molecule to the reaction product to provide a complex; and

21 2) determining the amount of bound biological 22 material by detecting the detectable molecule.

Preferably the detectable molecule is a monoclonal 1 antibody (MAb) specific to the biological compound. 2 Preferably the solid support is a microtitre or a 3 treated glass slide. 4 5 Preferably the method of assaying pyrrole-containing 6 biological compounds is Method 2 described in part b) 7 Method 2 involves the following steps: 8 9 contacting the biological compound with an 1) 10 optionally labelled derivatizing agent in 11 solution to form a reaction product therewith 12 (preferably via covalent attachment thereto) 13 followed by exposure to a binding agent bound to 14 a solid support, said binding agent being 15 specific to the biological compound in the 16 reaction product and 17 18 determining the amount of bound biological 2) 19 compound by determining the amount of labelled 20 derivatizing agent bound to the solid support. 21 22 Preferably the derivatizing agent is biotinylated 23 Ehrlich's reagent. Preferably the solution 24 containing the reaction product is neutralised prior 25 to contact with the bound binding agent. Preferably 26 the bound MAb is bound to a solid support, suitably a 27 microtitre plate or a treated glass slide. 28

Preferably the derivatizing agent is labelled with a 1 labeling molecule, suitably a radio-labelled, 2 fluorescent label, enzyme label or the like. 3 Preferably the amount of bound biological compound is 4 determined by detecting the amount of labelled 5 derivatizing agent bound on the solid support. 6 7 Method 2 takes into account the fact that relatively 8 strong acid conditions are required for the reaction 9 of derivatizing reagents with pyrroles. Thus, most 10 non-covalent interactions, such as antibody-antigen 11 complexes, would be disrupted under these conditions. 12 To overcome this problem, pyrrolic units in the 13 biological sample are targeted in Method 2 by 14 reaction in solution with derivatizing agent to form 15 a reaction product, preferably via covalent 16 attachment thereto followed by capture of the 17 reaction product on a surface coated with specific 18 antibodies. 19 20 Preferably, the method of assaying pyrrole-containing 21 biological compounds is Method 3, described in part 22 a) above. Method 3 involves the following steps: 23 24 contacting a biological compound with a 25 1) derivatizing agent in solution to form a 26 reaction product wherein the derivatizing agent 27 comprises a first partner of a strong binding 28

pair.

contacting the reaction product with a solid 2) 1 support having a second partner of the strong 2 binding pair on its surface, to form a bound 3 complex with the reaction product; 4 contacting the bound complex with a detectable 5 3) molecule; 6 determining the amount of bound biological 7 4) compound by detecting the amount of detectable 8 molecule bound to the solid support. 9 10 Preferably the derivatizing agent is a p-11 dimethylaminobenzaldehyde derivative, and in bound 12 form has the following structure: 13

$$\begin{array}{c}
H \downarrow O \\
R^{1} \downarrow N \\
R^{2} \downarrow Q \\
R^{4} \downarrow X:Y
\end{array}$$

14

wherein  $R^1$  is an alkyl group,  $R^2$  is an alkyl group,  $R^4$ 15 is a heteroalkyl group, X is a first partner of a 16 strong binding pair and Y is a solid support having a 17 second partner of a strong binding pair on its 18 surface. 19 20

Preferably the solution containing the reaction 21 product is neutralized prior to contact with the 22 solid support. 23

1	In one embodiment the first partner of the strong
2	binding pair is from avidin and the second partner of
3	the strong binding pair is from biotin.
4	Alternatively the first partner of the strong binding
5	pair is from biotin and the second partner of the
6	strong binding pair is from avidin. In a second
7	embodiment the first partner of the strong binding
8	pair is from biotin and the second partner of the
9	strong binding pair is from streptavidin.
10	Alternatively the first partner of the strong binding
11	pair is from streptavidin and the second partner of
12	the strong binding pair is from biotin.
13	
14	Preferably the detectable molecule is a monoclonal
15	antibody specific to the biological compound moiety
16	of the complex. Suitably the solid support is a
17	microtitre plate or a treated glass slide.
18	
19	The present invention also provides a method of
20	purifying an antigen, said method comprising;
21	
22	<ol> <li>contacting a pyrrole-containing biological</li> </ol>
23	compound with one of;
24	a) an optionally labelled derivatizing agent
25	(bound or able to bind to a solid support)
26	wherein the dirivatizing agent forms a
27	reaction product with the biological
28	compound (preferably via covalent
29	attachment thereto) followed by exposure to

1	a detectable molecule which forms a complex
2	with the reaction product; or
3	<ul><li>b) an optionally labelled derivatizing agent,</li></ul>
4	not bound to a solid support, wherein the
5	derivatizing agent forms a reaction product
6	with the biological compound (preferably
7	via covalent attachment thereto), followed
8	by exposure to a binding agent bound to a
9	solid support wherein the binding agent is
10	specific to a biological compound in the
11	reaction product; or
12	<ul> <li>a binding agent bound to a solid support,</li> </ul>
13	said binding agent being specific to the
14	biological compound, and forming a complex
15	therewith, followed by exposure to an
16	optionally labelled, derivatizing agent,
17	which forms a reaction product with the
18	biological compound moiety of said complex
19	(preferably via covalent attachment
20	thereto); and
21	2) eluting the biological compound from the solid
22	support.
23	
24	This method allows easy preparation of an antigen,
25	which can then be used in screening for an antigen
26	detection agent, for example antibody.
27	
28	Preferably the derivatizing agent for use in the
29	method of purifying an antigen is of the following
30	structure in bound form:

1 2 wherein  $R^1$  is an alkyl group,  $R^2$  is an alkyl group, A

3 is a linking group and B is a solid support.

4

5 Preferably the labeled derivatizing agent has the

6 following structure in bound form:

$$\begin{array}{c}
H \downarrow O \\
R^{1} \downarrow N \downarrow R^{2} \downarrow Q \\
R^{4} \downarrow X:Y
\end{array}$$

7 wherein  $R^1$  is an alkyl group,  $R^2$  is an alkyl group,  $R^4$ 

8 is a heteroalkyl group, X is a first partner of a

9 strong binding pair and Y is a solid support having a

10 second partner of a strong binding pair on its

11 surface.

12

13 Preferably the detectable molecule is a monoclonal

14 antibody specific to the biological compound.

15

16 Optionally the derivatizing agent is labelled with a

17 radio-label, fluorescent label, enzyme label or the

18 like.

1 The present invention also provides compounds for use

2 in the method of assaying pyrrole-containing

3 biological compounds.

4

5 In one case, the compound is of the following

6 structure:

7

8 wherein  $R^1$  is an alkyl group,  $R^2$  is an alkyl group, A

9 is a linking group and B is a solid support.

10

11 More preferably the labeled derivatizing agent has

12 the following structure:

wherein R1 is a straight-chain alkyl group containing 1 to 10 carbon atoms,  $R^2$  is a straight-chain alkyl group containing 1 to 10 carbon atoms, and  $R^4$  is a 3 straight-chain heteroalkyl group containing 2 to 10 4 carbon atoms and at least 2 heteroatoms. 5 6 BRIEF DESCRIPTION OF THE DRAWINGS 7 8 FIGS. 1a-1i show mass spectrometry spectra of pyrrole crosslink-containing peptides. 10 11 FIG. 2 schematically represents Methods 1, 2 and 3. 12 13 FIG. 3 shows the difference of pyrrole capture of 14 bone peptides at different dilutions. 15 16 FIG. 4 shows pyrrole capture at different dilution of 17 biological sample using detection antibodies specific 18 for isoaspartyl telopeptides. 19 20 FIG. 5 shows pyrrole capture assay for digested and 21 immobilized collagen-containing tissues. 22 23 FIG. 6 shows the results for a serial dilution of 24 biotin-ER reacted bone digest or a streptavidin 25 coated plate detected with NTP monoclonal antibody. 26

DESCRIPTION OF THE SPECIFIC EMBODIMENTS 1 2 Introduction 3 The present invention provides methods of assaying pyrrole-containing biological compounds and chemical 5 compositions that can be used in those methods. 6 Method 1 of the present invention, a biological 7 sample, that may have been processed, is contacted 8 with a solid support bound or able to bind 9 derivatizing agent. Pyrrolic units in the biological 10 sample react with the derivatizing agent, thereby 11 immobilizing components containing the pyrroles on 12 the solid support. The reacted solid support is 13 contacted with a detectable molecule, such as a MAb, 14 which interacts with a portion of the immobilized 15 biological material. Detection of the detectable 16 molecule on the solid support indicates that the 17 biological material contains pyrrolic units. 18 19 In Method 2 of the present invention an optionally 20 processed biological sample is contacted with a non-21 bound, optionally labeled derivatizing agent in 22 solution. The derivatizing agent is suitably labelled 23 with a radio-label, fluorescent label, enzyme label 24 or the like. The derivatizing agent reacts with the 25 pyrrolic units in the biological sample to form a 26 reaction product wherein the reaction product 27 comprises the covalent attachment of the derivatizing 28 agent and the pyrollic units in the biological 29

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compound. The solution containing the reaction
1
    product is neutralised.
2
3
    The reaction product may be contacted with a solid
4
    support bound MAb specific to the biological sample.
5
    The MAb reacts with the reaction product to form a
б
    complex immobilized on the solid support. Detection
    of the labeled molecule on the solid support
8
    indicates that the biological material contains
9
10
    pyrrolic units.
11
    In method 3 of the present invention, an optionally
12
    processed biological compound is contacted with a
13
    derivatizing agent, wherein the derivatizing agent
14
    comprises a first binding partner of a strong binding
15
    pair, suitably from biotin. The derivatizing agent
16
    is in solution. Pyrrolic units in the biological
17
    compound react with the derivatizing agent to form a
18
    reaction complex. The solution containing the
19
    reaction product is neutralised prior to contact with
20
    a solid support coated with a second binding partner
21
    of the strong binding pair, to form a bound complex
22
    with the reaction product. Suitably the second
23
    binding partner is from streptavidin. The solid
24
     support is then contacted with a detectable molecule,
25
    preferably a MAb specific to the biological compound
26
     moiety of said complex. The amount of bound
27
     biological compound is determined.
28
29
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FIG. 2 schematically illustrates Methods 1, 2 and 3. 30

## Definitions 1 "Alkyl group" refers to a straight-chain, branched or 2 cyclic group containing a carbon backbone and 3 hydrogen. Examples of straight-chain alkyl groups 4 include methyl, ethyl, propyl, butyl, pentyl and 5 hexyl. Examples of branched alkyl groups include i-6 propyl, sec-butyl and t-butyl. Examples of cyclic 7 alkyl groups include cyclobutyl, cyclopentyl and 8 cyclohexyl. The "alkyl" group also refers to 9 alkylene groups. 10 11 Alkyl groups are substituted or unsubstituted. 12 substituted alkyl group, a hydrogen on the carbon 13 backbone is replaced by a different type of atom 14 (e.g., oxygen, nitrogen, sulfur, halogen). 15 instance, 2-hydroxyethyl is an ethyl group where one 16 of the hydrogens is replaced by an OH group; 2-17 chloropropyl is a propyl group where one of the 18 hydrogens is replaced by a Cl group. 19 20 "Heteroalkyl group" refers to a straight-chain, 21 branched or cyclic group containing a carbon-22 heteroatom backbone and hydrogen. Heteroatoms 23 include, without limitation, oxygen, nitrogen and 24 The following groups are examples of sulfur. 25 heteroalkyl groups: $-CH_2OCH_2CH_3$ , $-NH(CH_2)_5NH-$ and 26 $-NH(CH_2)_2SS(CH_2)_2NHC(O)(CH_2)_5NH-$ . As with alkyl 27 groups, heteroalkyl groups are substituted or 28 unsubstituted. 29

"Leaving group" refers to a chemical group that is 1 capable of being displaced in a nucleophilic 2 substitution reaction. Examples of leaving groups 3 include -Cl, -Br, -OC(O)CH3 and -SPh. 4 5 "Linking group" refers to a chemical group that 6 connects one chemical group to another. 7 instance, in the compound  $CH_3C(O)-NH(CH_2)_5NH-CH_3$ , the 8 group -NH(CH<sub>2</sub>)<sub>5</sub>NH- is a linking group between CH<sub>3</sub>C(O)-9 and -CH3. 10 11 Types Of Biological Materials To Be Examined 12 13 The present method is used to determine the presence 14 of pyrrolic units in biological materials, including 15 pyrrolic crosslinks in collagen extracts. For some 16 time, researchers have proposed that pyrrolic 17 components exist in collagen. See Scott et al., 18 Biosci. Rep. (1981) 1:611-618; see also Kuypers et 19 al., Biochem. J. (1992) 283:129-136. Only indirect 20 support for the proposal has been available, however, 21 as the isolation and characterization of collagen 22 derived pyrrolic crosslinks has proven difficult. 23 24 Experimental results presented herein provide direct 25 confirmation of pyrrolic crosslinks in collagen. 26 Examples 4 and 5. A series of peptides from human 27 bone collagen enzyme digests were isolated using a

solid support bound p-aminobenzaldehyde, indicating

28

- 1 the presence of pyrrolic units in the collagen.
- 2 Analysis of the isolated peptides using mass
- 3 spectrometry showed that a relatively large number of
- 4 the peptides possessed masses extremely close to the
- 5 theoretic masses of complexes derivatized at
- 6 predominantly the N-telopeptide sites of collagen.

- 8 Pyrrolic crosslinks are particularly prevalent in
- 9 bone collagen where they result from the natural
- 10 maturation process of the tissue. During resorption
- 11 of bone by osteoclasts, fragments of collagen
- 12 crosslinked by pyrroles are released into the
- 13 circulation. Their concentration in various
- 14 biological fluids provides an indication of the rates
- 15 of bone degradation. Increased bone resorption rates
- 16 are associated with a number of diseases, including,
- 17 for example, the following: osteoporosis, osteo- and
- 18 rheumatoid arthritis, and diseases involving
- 19 abnormalities of vitamin D or parathyroid hormone
- 20 such as osteomalacia and hyperparathyroidism. By
- 21 detecting pyrrolic crosslinks using the present
- 22 invention, therefore, one is able to characterize and
- 23 monitor such diseases.

- 25 Another example of biological materials that can be
- 26 assayed using the present invention is the
- 27 isolevuglandins (e.g., levuglandin  $E_2$ ).
- 28 Isolevuglandins are formed through free radical-
- 29 mediated oxidation of polyunsaturated fatty acid
- 30 esters in low-density lipoproteins. These compounds

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react with various proteins to produce pyrroles in
    vivo. See Brame et al., J. Biol. Chem. (1999)
2
    274:13139-13146; see also Salomon et al., J. Biol.
3
    Chem. (1999) 274:20271-20280.
4
5
    Free radical-mediated oxidation has been implicated
6
    in a wide variety of human diseases, including
7
    atherosclerosis, cancer and neurodegenerative
8
    diseases. See B. Halliwell and J. Gutteridge,
9
    Methods Enzymol. (1990) 186:1-85. Specifically, the
10
    oxidative modification of low density lipoproteins is
11
    a key step in atherosclerosis etiology.
12
    detection of isolevuglandin derived pyrroles
13
    accordingly provides a method for diagnosing and
14
    monitoring atherosclerosis.
15
16
    Proteins modified by non-enzymatic glycosylation
17
    reactions constitute a third example of a biological
18
    material that can be assayed using the present
19
    invention. Threose, primarily derived from the
20
    breakdown of ascorbate (vitamin C), represents one
21
                                 It is particularly
     instance of this reaction.
22
    reactive with lysine residues in proteins and forms
23
    pyrrolic structures (e.g., formyl threosyl pyrrole)
24
     as a result. See R. Nagaraj and V. Monnier, Biochem.
25
     Biophys. Acta (1995) 1253:75-84.
26
27
     Detecting formyl threosyl pyrrole is specifically
28
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useful for monitoring patients with diabetes.

also an example of an advanced glycation end-product

29

(AGE). AGEs are associated, for example, with 1 abnormal neurofibrillar structures in Alzheimer's 2 disease, and the presence of increases AGEs in 3 lipoproteins appears to accelerate the oxidative 4 reactions leading to atherosclerosis. Therefore, the detection of formyl threosyl pyrrole provides a method for diagnosing and monitoring those diseases as well. 8 . 9 Methods Of Processing Biological Materials 11 Subject biological materials assayed using the 12 present method may be unprocessed (e.g., urine, serum 13 or plasma) or processed. A primary goal of 14 processing is the solubilization of the sample. 15 16 Where the biological material is a tissue, it is 17 usually de-fatted by two brief extractions (e.g., 15 18 min.) with acetone or chloroform: methanol (2:1 v/v). 19 Mineralized tissues are, for example, powdered 20 underliquid nitrogen and subsequently demineralized 21 using extraction with 0.5 M EDTA at pH 7.5 for 72-96 22 hours at 4 °C. Connective tissue samples are 23 typically denatured by heating the sample in saline 24 at pH 7.4 for 30 min at 70 °C. 25 26 Sample solubilization typically involves the use of 27 proteases rather than chemical hydrolysis, as 28

pyrroles exhibit chemical instability under certain

conditions. Where proteases are used, a sample is

29

- treated with a suitable proteolytic enzyme (e.g., 1
- trypsin) at a suitable temperature (e.g., 37 °C). 2
- Examples of other enzymes one can use to solubilize a 3
- biological material include chymotrypsin, pronase, 4
- pepsin, proteinase K and members of the cathepsin 5
- family (B, L, N or K). For any chosen enzyme, one of 6
- ordinary skill can readily determine a suitable 7
- reaction buffer pH and temperature. 8

## Derivatizing Agents 10

11

- The deritivizing agents used in the present assay are 12
- p-amino benzaldehyde derivatives used in the present 13
- assay are of the structures  ${\bf 4}$  and  ${\bf 5}$ .  ${\bf R}^{\bf 1}$  in the 14
- structures is an alkyl group; R2 is an alkyl group; 15

16

- R3 is a hydroxyl group or leaving group; and, R4 is a 17
- heteroalkyl group. 18

- The substituent R1 is preferably a straight-chain 20
- alkyl group containing 1 to 10 carbon atoms. 21
- more preferably a straight-chain alkyl group 22

containing 1 to 5 carbon atoms. Most preferably, R1 1 contains 1 carbon atom (i.e.,  $-CH_3$ ). 2 3 The substituent R2 is preferably a straight-chain 4 alkylene group containing 1 to 10 carbon atoms. 5 is more preferably a straight-chain alkylene group containing 1 to 5 carbon atoms. Most preferably  ${\ensuremath{\mbox{R}}}^2$ 7 contains 2 carbon atoms (i.e.,  $-CH_2CH_2-$ ). 8 The substituent  $R^3$  is preferably -OH, -OR $^5$  (where  $R^5$ 9 is a straight chain alkyl such as methyl), -Cl or 10 SR<sup>5</sup>. It is more preferably -OH or -OR<sup>5</sup>. Most 11 preferably R3 is -OH. 12 13 The substituent R4 is preferably a straight-chain 14 heteroalkyl group containing 2 to 10 carbon atoms and 15 at least 2 heteroatoms. It is more preferably a 16 straight-chain heteroalkyl group containing 4 to 10 17 carbon atoms and at least 2 nitrogen atoms. 18 preferably R4 is-NHCH2CH2CH2CH2CH2CH2NH- or 19  $-\mathrm{NHCH_2CH_2SSCH_2CH_2NHC}\left(\mathrm{O}\right) - \mathrm{CH_2CH_2CH_2CH_2CH_2N} -.$ 20 21 Examples of three preferred derivatizing agents are 22 p-amino benzaldehyde derivatives are shown as 23 compounds 6, 7 and 8: 24

$$\begin{array}{c} H \longrightarrow O \\ \\ H_3C \longrightarrow NH(CH_2)_2SS(CH_2)_2NHC(O)(CH_2)_5NH \longrightarrow O \\ \\ R \longrightarrow NH \longrightarrow NH \end{array}$$

## Modes Of Attachment To A Solid Support 2

3

- The derivatizing agent is attached to the solid 4
- support through either a covalent bond or a 5
- noncovalent interaction. A derivatizing agent in 6
- bound form attached to solid support through a 7
- covalent bond is represented by compound 9; a 8
- derivitizing agent in bound form attached to a solid 9
- support through a noncovalent interaction is 10
- represented by compound 10: 11

12

- The substituents of compound 9 are defined as
- follows: R1 is an alkyl group; R2 is an alkyl group; 14
- A is a linking group and B is a solid support. 15
- Preferably,  $R^1$  and  $R^2$  are alkyl groups containing 1 16
- to 10 carbon atoms and A is a heteroalkyl group. 17

- 1 More preferably, R1 and R2 are alkyl groups
- 2 containing 1 to 5 carbon atoms and A is a heteroalkyl
- 3 group comprising at least 1 nitrogen atom. Most
- 4 preferably,  $R^1$  is  $-CH_3$  and  $R^2$  is  $-CH_2CH_2-$ . The
- 5 substituents of compound 10 are defined as follows:
- 6  $R^1$  is an alkyl group;  $R^2$  is an alkyl group;  $R^4$  is a
- 7 heteroalkyl group; X is a first partner of a strong
- 8 binding pair and Y is a solid support having a second
- 9 partner of a strong binding pair on its surface.
- 10 Preferably, R<sup>1</sup> and R<sup>2</sup> are alkyl groups containing 1
- 11 to 10 carbon atoms and R4 is a straight-chain
- 12 heteroalkyl group containing 2 to 10 carbon atoms and
- 13 at least 2 heteroatoms. More preferably,  $R^1$  and  $R^2$
- 14 are alkyl groups containing 1 to 5 carbon atoms,  $exttt{R}^4$
- 15 is  $-NH(CH_2)_5NH-$  or  $-NH(CH_2)_2SS(CH_2)_2NHC(O)(CH_2)_5NH-.$
- 16 Most preferably,  $R^1$  is  $-CH_3$  and  $R^2$  is  $-CH_2CH_2-$ .

- 18 Where a covalent bond is used for attachment, a
- 19 surface is typically derivatized to afford a reactive
- 20 functional group such as an alcohol or amine. For
- 21 instance, compound  $\mathbf{6}$  is coupled to a Nunc Covalink<sup>TM</sup>
- 22 plate, available from Nalge Nunc International,
- 23 through the formation of an amide bond with a C8-
- 24 primary amine. See www.nalgenunc.com. A second
- 25 example of a suitable solid support is a DNA-BIND<sup>TM</sup>
- 26 surface, available from Corning. See
- 27 www.scienceproducts.corning.com. One reacts a
- 28 bifunctional compound, such as 1,5-diaminopentane,
- 29 with the surface to provide available amine groups

- 1 for covalent attachment. A compound such as 6, which
- 2 contains a carboxylic acid, is coupled to the surface
- 3 groups through the formation of an amide bond. A
- 4 third example of a solid support is a glass
- 5 substrate. A glass slide is treated with
- 6 aminopropyl-triethoxysilane to provide a glass
- 7 substrate containing a reactive amine across its
- 8 surface. See U.S. 5,919,523. The derivatized slide
- 9 is reacted with compound such as 6 in the presence of
- 10 a suitable reagent that induces amide bond formation.
- 11 Where a noncovalent interaction is used for
- 12 attachment, a compound containing one partner of a
- 13 strong binding pair is adhered or bonded to the solid
- 14 support. The other partner of the pair is covalently
- 15 attached to a derivatizing agent to form a conjugate.
- 16 When the conjugate is contacted with the solid
- 17 support, a strong interaction (e.g., one or more
- 18 hydrogen bonds) immobilizes the conjugate on the
- 19 support.
- 20
- 21 An example of a strong binding pair is a
- 22 biotin:avidin complex. (A biotin:streptavidin
- 23 complex is another example.) Typically, a support
- 24 surface is derivatized to include biotin or avidin.
- 25 Avidin coated polystyrene plates (i.e., Reacti-Bind<sup>TM</sup>
- 26 NeutrAvidin<sup>TM</sup> coated plates) are available, for
- 27 instance, from Pierce. See www.piercenet.com. The
- 28 avidin coated plate is contacted with a biotin
- 29 containing p-aminobenzaldehyde derivative such as
- 30 compound 7. The resulting biotin-avidin complex

1	serves to attach compound 7 to the solid support
2	through noncovalent interactions.
3	
4	Examples Of Different Assay Formats
5	
6	The method of assaying pyrrole-containing biological
7	compounds is typically run in a multi-well plate
8	(e.g., 96-well plate), but other assay formats are
9	also used. The method is also performed using a
10	strip format, where a derivatizing agent is
11	immobilized on the strip surface. A third exemplary
12	format involves the use of a polymeric bead (e.g.,
13	polystyrene bead) on which a derivatizing agent is
14	immobilized. Yet another format involves the use of
15	micro-array or chip technology; use with surface
16	plasmon resonance technology.
17	
18	Contact Of Extract/Isolate With Detection Compound
19	
20	To perform a method of the present invention, a
21	biological fluid or processed biological material is
22	contacted with a solid support bound derivatizing
23	agent. The biological material may be solubilized in
24	a suitable solvent to form a solution prior to the
25	contact. When a multi-well format is used, for
26	example, the solution and any additional elements
27	readily discernable to one of ordinary skill in the
28	art is added to one or more wells. For the strip
29	format, a strip is dipped into a solution containing
30	the biological material; and, for the bead format, a

vial or tube is used to mix the beads and the 1 solution. 2 3 Regardless of assay format, contact between a 4 pyrrole-containing biological material and the 5 support bound derivatizing agent induces a coupling reaction. The result of the reaction is a covalent 7 bond between the biological material and the 8 derivatizing agent. This serves to immobilize the 9 pyrrole-containing biological material on the solid 10 support. 11 12 When desired, the solid support bound biological 13 material is washed with at least one suitable solvent 14 to remove impurities from the reaction medium. 15 solid support is typically dried after a washing 16 step. A variety of drying techniques are used, 17 including air drying, drying under reduced pressure 18 and thermal drying. 19 20 Methods Of Detection Using A Detectable Molecule 21 22 In a method of the present invention, the immobilized 23 material is contacted with a detectable molecule. 24 The detectable molecule specifically binds to a 25 portion of a targeted biological material. 26 material on the solid support is not the targeted 27 material, the detectable molecule will not bind to it 28 with high affinity. 29

- The detectable molecule can bind to the targeted 1 biological material through either covalent or 2 noncovalent bonds. Typically, the detectable 3 molecule is a polyclonal, monoclonal or phase 4 library-derived antibody that binds to the biological 5 material through noncovalent bonds. Preferably, it 6 is a monoclonal antibody. 7 8 The detectable molecule is typically detectable in 9 one of three ways: 1) it contains functionality one 10 can observe; 2) it induces a chemical reaction that 11 . produces an observable product; or 3) it interacts 12 with a second molecule that either contains 13 functionality one can observe or induces a chemical 14 reaction that produces an observable product. 15 Functionality one can observe includes chemical 16 groups that exhibit a measurable effect upon 17 stimulation. For instance, the following chemical 18 groups exhibit such an effect: a chemical group that 19 absorbs light at a certain wavelength (a chromophore) 20 and a chemical group that fluoresces upon exposure to 21 a particular wavelength of light. A chemical 22 reaction that produces an observable product 23 includes, for example, a reaction producing a 24 fluorescent compound, a luminescent compound or a 25 chromophoric compound. 26 27 Where the targeted biological material is collagen
- 28
- derived pyrrole crosslinks, an example of a 29
- detectable molecule is a monoclonal antibody (NTP) 30

raised against a synthetic octapeptide comprising 1 part of the sequence of the  $\alpha 2$ (I) N-terminal 2 telopeptide. The NTP antibody is contacted with the 3 immobilized biological material. A secondary 4 antibody (goat anti-mouse IgG-peroxidase conjugate) 5 is introduced; which interacts with a portion of the 6 NTP antibody. Upon addition of 3,3',5,5'-7 tetramethyl-benzidine dihydrochloride and hydrogen 8 peroxide, a chromophoric compound exhibiting an 9 absorbance at 450 nm is produced. See Example 6. 10 11 Contact of Extract/Isolate with Detection Compound 12 To perform Method 2 or 3 of the present invention, a 13 biological fluid or processed biological material is 14 contacted with a labeled derivatizing agent in 15 The derivatizing agent is labeled with a solution. 16 labeling molecule. Any suitable solvent as known by 17 a person skilled in the art may be used. A coupling 18 reaction between pyrrole-containing biological 19 material results in a reaction product comprising the 20 derivatizing agent covalently bonded to any pyrrole-21 containing biological material. 22 23 Methods of Detection Using a MAb 24 In Method 2 of the present invention the reaction 25 product is immobilised by contact of the solution 26 with a MAb bound on a solid support. 27

29 30

- 1 Example 1: Preparation of compound 6.
- 2 N-Methyl-N-cyanoethyl-4-amino benzaldehyde (available
- 3 from Enterwin Chemicals, China or Sigma-Aldrich, USA)
- 4 (150 mg) was dissolved in 7.5 M NaOH, 6%  $H_2O_2$  (5 ml)
- 5 and refluxed for 2 hours. The hydrolysate was
- 6 acidified by addition of concentrated HCl, dried
- 7 under vacuum and redissolved in ethanol (1.5 ml). An
- 8 aliquot of the solution (1 ml) was added to 0.2 M
- 9 NaOH (1 ml) and applied to an anion exchange column
- 10 (Bio-Rad AG 1-X8; 2 ml, pretreated with 2 M HCl, 2 M
- 11 NaOH and equilibrated with water). The column was
- 12 washed with water (12 ml) before elution of the bound
- 13 material with 2 M HCl. The eluent was dried under
- 14 vacuum and the residue resuspended in water (1 ml).
- 15 A small amount of residue (soluble in ethanol but
- 16 containing no compound 6) was removed after which the
- 17 aqueous fraction was dried under vacuum (yielding 7
- 18 mg of material) and redissolved in 0.1%
- 19 trifluoroacetic acid (1 ml). Aliquots (100  $\mu$ l) of
- 20 the material was chromatographed on a Waters RCM
- 21 Prep-Pak® C<sub>18</sub> column (25 mm x 100 mm, 10 μm) pumped
- 22 at 4 ml/min. The buffers used were 0.1% TFA (buffer
- 23 A) and 70% acetonitrile, 0.1% TFA (buffer B) with a
- 24 gradient of 5 minutes at 5% B followed by a linear
- 25 increase to 70% B over 35 minutes. Monitoring at 330
- 26 nm showed a single major peak which eluted at 28.3
- 27 min. Fractions corresponding to the peak were pooled
- 28 and dried under vacuum (yield = 3 mg). Analysis of
- 29 the material by electrospray mass-spectrometry in
- 30 negative-ion mode using a MAT 900 mass spectrometer

(Finnigan MAT, Bremen, Germany) revealed the major 1 ion as [M-H] = 206.2 which corresponds to the 2 expected value for N-methyl-N-propionic acid-4-amino 3 benzaldehyde  $M_r$  207.2. 4 5 Example 2: Preparation of compound 7. 6 Compound 6 (3 mg) was dissolved in water (3 ml) and 7 biotin-pentyl amine (30 mg; Pierce) was added. 8 solution of 1-ethyl-3-(3-dimethylamino-9 propyl)carbodiimide/N-hydroxysuccinimide (0.035 10 M/0.028 M respectively; 3 ml) was added and heated to 11 50 °C for 4 h. The resulting solution was dried 12 under vacuum and chromatographed using the 13 preparative RCM Prep-Pak® column described in 14 Example 1. The gradient applied was 20% B for 5 min 15 followed by a linear increase to 60% B over 30 min. 16 Two major components were detected, one eluting at 15 17 min. (unreacted acid) and one eluting at 18 min. 18 component eluting at 18 min was analyzed by positive-19 ion electrospray mass-spectrometry and showed [M+H] 20 of 518.7 and [M+Na] of 540.6. These values 21 corresponded to the calculated  $\ensuremath{\text{M}_{\text{r}}}$  of compound 7 of 22 517.7. Compound 7 reacted with pyrrole carboxylic 23 acid in 4 M HCl to give a characteristic pink color 24 absorbing at 573 nm. 25

- 27 Example 3: Preparation of compound 8.
- 28 Compound 6 (1 mg) was dissolved in 0.1 M MES buffer
- 29 pH 5 (1 ml) and a ten-fold molar excess of cystamine
- 30  $(H_2N(CH_2)_2SS(CH_2)_2NH_2)$  was added. The solution pH was

- 1 adjusted to 5 using HCl, and a solution of 1-ethyl-3-
- 2 (3-dimethylamino-propyl)-carbodiimide/N-
- 3 hydroxysuccinimide (0.035 M/0.028 M respectively; 1
- 4 ml) was added. The solution was heated to 50 °C for
- 5 4 h. The resulting aminated derivative was purified
- 6 by HPLC, eluting with 10 mM TFA and an acetonitrile
- 7 gradient (monitoring 330 nm). Biotinylation of the
- 8 aminated derivative was performed using succinimide-
- 9 LC-biotin (Pierce) according to the manufacturer's
- 10 instructions and again purified by HPLC. Structure 8
- 11 was confirmed by MALDI-TOF mass spectrometry.

- 13 Example 4: Reaction of compound 7 with a bone
- 14 digest.
- 15 De-fatted human bone (7 g) was powdered in a Spex
- 16 freezer-mill in liquid nitrogen. The resultant
- 17 powder was decalcified by 3 x 2-day extractions in
- 18 0.5 M EDTA, pH 8 at 4 °C, washed with water and
- 19 lyophilized. The decalcified bone powder (1.1 g) was
- 20 suspended in 0.1 M citrate buffer, pH 5, heated to
- 21 70°C for 1 hour to denature the triple-helical
- 22 structure and allowed to cool to 45 °C. Papain (100
- 23 U) was added, and the digest was incubated for 4
- 24 hours. The pH of the digest was adjusted to 7.4 by
- 25 the addition of 1 M Tris, and the temperature was
- 26 lowered to 37 °C for an overnight digestion with
- 27 protease type X (100 U). The completed digest
- 28 (estimated as 110  $\mu M$  collagen by total pyridinium

crosslink content) was frozen, lyophilized and 1 suspended in water (7 ml). 2 3 After the addition of compound 7 (50  $\mu g$ ) to the bone 4 digest (500  $\mu$ l), the mixture was acidified by the 5 addition of 12 M HCl (250  $\mu$ l). During incubation for 6 30 min at room temperature, the solution turned 7 cherry-pink in color, and spectrometry showed the 8 presence of an absorption maximum at 571.7 nm 9 (characteristic of product from reaction of 4-10 dimethylamino benzaldehyde with pyrrole). The acid 11 was neutralized by the addition of 12 M NaOH (approx. 12 220  $\mu$ l) followed by 40 mM phosphate buffer (20 ml). 13 14 Example 5: Isolation of conjugation product between 15 compound 7 and pyrrolic peptides. 16 A monomeric avidin column (5 ml) was prepared 17 according to manufacturer's (Pierce) instructions. 18 The reacted bone digest of Example 4 at neutral pH 19 was added slowly to the column, which was then washed 20 with 6 column volumes of PBS followed by 1 column 21 volume of water. The biotinylated material was 22 eluted at about 1 ml/min with 1 M acetic acid 23 adjusted to pH 2.5 with ammonia, and 8 fractions (5 24 ml) were collected. 25 26 Estimation of biotinylated compounds by competitive 27 ELISA. In order to assess the efficiency of the 28 monomeric avidin column (Example 5), a competitive

ELISA was developed. Immulon 4 immunoassay plates 1 were coated with streptavidin (25 nM) in PBS for 2 2 hours at 37 °C. Samples or standards in PBS 0.1% 3 Tween, 0.5% fat-free milk powder (FFMP; 110  $\mu$ l) were 4 added to biotinylated peroxidase (Sigma; 10 ng/ml; 5 110  $\mu$ l) in PBS Tween, 0.5% FFMP in a U-bottomed 96-6 well plate. The mixed samples were transferred to 7 the washed, streptavidin-coated plate and incubated 8 for 90 min at 37 °C. After washing the plate 3 times 9 with PBS/0.1% Tween, the peroxidase substrate (200 10  $\mu$ l) tetramethyl-benzidine dihydrochloride (TMB) was 11 added (0.1 mg/ml) in 0.05 M citrate/phosphate buffer 12 pH 5, 0.012% v/v hydrogen peroxide. The reaction was 13 stopped by the addition of 3 M sulphuric acid (50  $\mu$ l) 14 after 15 min. 15 16 Analysis of isolated material by HPLC. Material 17 eluted from the avidin column was reduced in volume 18 (100  $\mu$ l) and chromatographed on a reversed phase HPLC. 19 column (4.6 x 100 mm;  $C_{18}$ ; particle size 3  $\mu m$ ). The 20 column was equilibrated with 0.1% TFA (buffer A), and 21 peptides were eluted over 35 min with linear 22 gradients formed with 70% acetonitrile, 0.1% TFA 23 (buffer B). The eluent was monitored at 214 nm, 280 24 nm and at 330 nm. Each fraction from the HPLC was 25 dried and redissolved in water (2  $\mu$ l). An aliquot (1 26  $\mu$ l) was mixed with  $\alpha$ -cyano-4-hydroxy-cinnamic acid (1 27  $\mu$ l of a 10 mg/ml solution in 70% acetonitrile 0.1% 28

TFA), dried onto a sample plate and analyzed by

MALDI-TOF mass spectrometry (Voyager-DE; Applied 1 Biosystems) calibrated externally using bradykinin. 2 3 The MALDI-TOF mass spectrometry spectra of each 4 fraction is shown in FIG. 1. As there were 5 insufficient quantities of many of the smaller 6 peptides to obtain amino acid composition data, some 7 ambiguities in their structural assignments did 8 In particular, the mass difference between 9 Glu and Ile/Leu is equivalent to an additional 10 hydroxyl group and, for the isolated peptide with  $M_{\mathtt{r}}$ 11 = 1086 (FIG. 1a), the ambiguity is due to the 12 possible presence of a hydroxylated crosslink. 13 this peptide may contain Gly and Glu (from either the 14 C- or N-telopeptides of the lpha 1 chain) or, for a 15 hydroxylated crosslink, a Gly residue linked with 16 either Ile (from the  $\alpha 1$  helix) or a leucine (from the 17  $\alpha$ 2 helix). Even where the amino acid composition is 18 known, the precise location of the residues may not 19 be clear, as in the case of the peptide with  $M_{\rm r}$  = 957 20 (FIG. 1a) containing the biotinylated pyrrole with a 21 single Gly residue. This residue is shown in a 22 helical position (which could be at the N- or C-23 terminal overlap sites) but could also be derived 24 from the  $\alpha 2$  (I) N-telopeptide: this peak may contain 25 a mixture of Gly-containing peptides from the 26 different locations. The Mr = 1029 peptides shown in 27 FIG. 1e and 1g could have the same alternatives of 28

glutamate or hydroxylated pyrrole-leucine/isoleucine.

- 1 The peaks corresponding to a loss of Gly (FIG. 1b, 2 lc) are probably losses due to the energy of the
- 3 laser-desorption rather than discrete peptides, but
- 4 these peaks provide additional evidence for the
- 5 peptide structures proposed. The structures of the
- 6 larger peptides shown in the other panels are
- 7 unambiguous.

- 9 Example 6: Detection of pyrrole crosslinks (Method 1)
- 10 The carboxyl-Ehrlich derivative was coupled to a Nunc
- 11 Covalink® plate via a C8-primary amine group. After
- 12 adding the derivative to the plate (250 pmole/well in
- 13 100  $\mu$ l MES buffer, pH 4.5) followed by 100  $\mu$ l of 1-
- 14 ethyl-3-(3-dimethylamino-propyl)carbodiimide/N-
- hydroxysuccinimide (0.035M / 0.028M respectively),
- 16 the plate was heated to 50°C and left overnight at
- 17 room temperature. The plate was aspirated and washed
- 18 with 4M HCl and 3 times with water. Each well
- 19 coupled the equivalent of 66 pmoles of the reagent
- 20 and the coupling was confirmed using HPLC.

- 22 Samples (110μ1), prepared in a separate plate, were
- 23 acidified by the addition of 8M HCl (110  $\mu$ l). The
- 24 acidified samples (200  $\mu$ l) were then added to the
- 25 Ehrlich reactive plate and agitated for 1 hour at
- 26 room temperature. The plate was aspirated and washed
- 27 3 times in 4 M HCl, 3 times in water and finally 3
- times in PBS/0.1% Tween; 10mM lysine, 0.5% fat-free
- 29 milk powder (assay buffer). The antibodies used were

- a monoclonal antibody (NTP) raised against the  $\alpha 2 \, (\text{I})$ 1 telopeptide (1:1000 dilution) or affinity-purified, 2 polyclonal antibodies raised against the isoaspartyl 3  $\alpha 2$ (I) telopeptide (1:250 dilution). After incubation 4 for 17 hours at 4 °C, the plate was washed 3 times 5 with PBS-Tween and incubated for 1 hour with 6 secondary antibodies, goat anti-mouse IgG-peroxidase 7 conjugate, used at a dilution of 1:4000. The plate was washed 3 times with PBS-Tween, and 200  $\mu$ l of 9 peroxidase substrate, 3,3',5,5'-tetramethyl-benzidine 10 dihydrochloride (TMB) is added (0.1 mg/ml) in 0.05 M 11. citrate/phosphate buffer, pH 5, containing 0.012% v/v 12 hydrogen peroxide. The reaction is stopped by the 13 addition of 3 M sulfuric acid (50  $\mu$ 1), and the 14 absorbance was measured at 450 nm using a Dynatech MR 15 7000 plate reader. 16 17 Using the pyrrole-capture assay, serial dilutions of 18 a bone digest (starting at ~1.0 nmole/well collagen) 19 reacted in the Ehrlich plate gave progressively 20 decreasing reactivity with NTP antibody (FIG. 3). 21 a fixed concentration (0.125 nmole/well) of pyrrole-22 crosslinked bone peptides on the plate, preincubation 23 of the NTP antibody with serial dilutions adolescent-24 human urine gave essentially complete inhibition of 25 colour development.
  - 26 27
  - 28 When pyrrole crosslink-containing peptides in urine
  - 29 from an adolescent were reacted with the plate, the

- 1 NTP antibody failed to detect any telopeptide (FIG.
- 2 4). A possible explanation for this is that the
- 3 large quantities of non-isomerised telopeptide found
- 4 in urine at this age may not be extensively
- 5 crosslinked. This is supported by the fact that the
- 6 polyclonal antibody raised against the isoaspartyl
- 7 rearranged peptide did show reactivity towards
- 8 captured peptides in urine from an older subject (30
- 9 years), see FIG. 4.

- 11 The specificity of the assay was demonstrated by
- 12 showing that peptides derived from cartilage and
- 13 skin, which have no pyrrolic crosslinks, gave very
- 14 little reaction in the assay compared to the bone
- 15 digest and a phorphobilinogen standard (FIG. 5).

16

- 17 Example 7: Detection of pyrrole-containing peptides
- 18 from enzyme digests of bone (Method 2)
- 19 A tryptic digest of demineralized human bone (0.5 ml
- 20 containing approximately 5 μM collagen) was reacted
- 21 with biotinylated Ehrlich's reagent (50  $\mu$ g; 0.1
- 22 μmoles) in 3MHCl for 30 min at room temperature. The
- 23 sample was neutralized by the addition of 2M NaOH and
- 24 diluted to 10 ml in phosphate buffered saline, pH 7.5
- 25 (PBS) containing 0.1% Tween 20. Serial (x2)
- 26 dilutions of this pre-reacted mixture were prepared
- 27 in PBS-Tween for addition to the detection plate.

- 29 The detection microtitre plate was coated with a
- 30 monoclonal antibody (NTP) recognizing an octapeptide

- 1 sequence containing the cross-linking region of the
- 2 N-telopeptide of collagen type I α2 chain. In order
- 3 to gain the appropriate orientation of the antibody,
- 4 the plate was initially coated (3 hours at room
- 5 temperature) with anti-mouse IgG (raised in donkey)
- 6 by adding to each well 0.2 ml of a solution
- 7 containing1 μg/ml protein in PBS. After washing 3
- 8 times with PBS-0.05% Tween 20, the NTP antibody (1
- 9  $\mu$ g/ml in PBS) was added and reaction allowed to
- 10 proceed for 1 hour at room temperature. The plate
- 11 was again washed 3 times with PBS-Tween.

- 13 Serial dilutions of the pre-reacted mixture were
- 14 added to the coated plate and incubated at room
- 15 temperature for 2 hours. The plate was washed 3
- 16 times with PBS-Tween and the biotin-pyrrole detected
- 17 by the addition of streptavidin-horseradish
- 18 peroxidase (Amersham plc, Little Chalfont, UK)
- 19 diluted 1:2000 in PBS-Tween. After 1 hour the plate
- 20 was washed 3 times in PBS-Tween and the colour
- 21 developed by the addition of 200  $\mu l$  of peroxidase
- 22 substrate, 3,3',5,5'-tetramethyl-benzidine
- 23 dihydrochloride (TMB) is added (0.1 mg/ml) in 0.05 M
- 24 citrate/phosphate buffer, pH 5, containing 0.012% v/v
- 25 hydrogen peroxide. The reaction is stopped by the
- 26 addition of 3 M sulfuric acid (50  $\mu$ l), and the
- 27 absorbance was measured at 450 nm using a Dynatech MR
- 28 7000 plate reader.

Example 8: Detection of pyrrole-containing peptides 1 from enzyme digests of bone (Method 3) Biotinylated Ehrlich's reagent was reacted with 2 tryptic peptides of human bone collagen as described 3 for Method 2. 5 For the detection plate, high-binding microtitre 6 plates (Immunlon 4) were coated with streptavidin (1 7  $\mu$ g/ml in PBS) by incubating for 3 hours at 37°C. 8 plates were washed 3 times with PBS-Tween and any 9 remaining binding sites were blocked by incubation at 10 room temperature for 1 hour with 3% bovine serum 11 The plate was again washed 3 12 albumin in PBS-Tween. Alternatively, ready coated 13 times with PBS-Tween. plates are available commercially from several 1.4 sources, such as Streptavidin-coated Combiplates from 15 16 Thermo Labsystems, Basingstoke, UK. 17 Serial dilutions of the pre-reacted mixture were 18 added to the streptavidin coated plate and incubated 19 at room temperature for 2 hours. The plate was 20 washed 3 times with PBS-Tween and, after the addition 21 of NTP monoclonal antibody (1:1000 dilution in PBS-22 Tween), the plate was incubated at 4°C for 18 hours. 23 The plate was washed 3 times with PBS-Tween and 24 incubated for 1 hour with secondary antibodies, goat 25 26 anti-mouse IgG-peroxidase conjugate, used at a dilution of 1:4000. After washing the plate 3 times 2'7 with PBS-Tween, colour development with TMB and 28 29

recording optical densities at 450 nm using the plate 1 reader were done as described previously. 2 3 Example 9: Preparation of pyrrole containing antigens 4 from bone collagen peptides 5 Peptides were prepared from powdered, decalcified 6 human bone by digestion with cathepsin K. 7 (10mg) was suspended in 1.0ml of 50mM sodium acetate 8 buffer, pH 5.0, containing 2mM EDTA and 2mM 9 dithiothreitol and, after the addition of 0.1mg 10 recombinant cathepsin K dissolved in 100µl PBS, 11 digestion was continued for 21 hours at 37°C with 12 gentle agitation. The digest was centrifuged 13 (13,000g) to remove any undigested tissue, and the 14 supernatant solution was desalted on a column (1.0 x 15 12cm) of Sephadex G25 equilibrated and eluted with 16 0.2M acetic acid. Pooled fractions containing the 17 bone peptides were lyophilised and reacted with 18 biotinylated, disulphide Ehrlich's reagent (compound 19 8; 0.1mg; 0.2µmoles) in 3M HC1 at room temperature 20 for 30 mins. The solution was neutralized by the 21 addition of 2 M NaOH and diluted to 10ml with PBS. 22 23 The bone digest Ehrlich conjugate was applied to a 24 5ml column of immobilized avidin (Pierce Chemical Co) 25 prepared according to the manufacturer's 26 instructions, and the column washed with PBS 27 containing 10 mM dithiothreitol and located by 28 monitoring the column effluent at 230nm. 29 fractions were dialysed against PBS to remove

- 1 reducing agent. This material was mixed with an
- 2 equal volume of adjuvant and used directly for
- 3 immunization of rabbits and mice.